Search of phenotype related candidate genes using Gene Ontology-based semantic similarity and protein interaction information: Application to Brugada syndrome.

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Abstract— This work presents a methodology for finding phenotype candidate genes starting from a set of known related genes. This is accomplished by automatically mining and organizing the available scientific literature using Gene Ontology-based semantic similarity. As a case study, *Brugada* syndrome related genes have been used as input in order to obtain a list of other possible candidate genes related with this disease. *Brugada* anomaly produces a typical alteration in the Electrocardiogram and carriers of the disease show an increased probability of sudden death. Results show a set of semantically coherent proteins that are shown to be related with synaptic transmission and muscle contraction physiological processes.

I. INTRODUCTION

Recent years have seen a drastic increase of the publicly available genetic and proteomic information. There are two main entry points for accessing gene and protein information. In the first place, UniProtKB[1], which is a collaboration between the European Bioinformatics Institute (EBI), the Swiss Institute of Bioinformatics (SIB) and the Protein Information Resource (PIR). Secondly, Entrez[2], which is a cross-database search tool offered by the National Center for Biotechnology Information (NCBI) from the US. This portal facilitates access to many available resources. In particular, Entrez Gene is a central source of information on genes, and transcripts. From all the publicly available sources of information on protein-protein interactions (PPI), three of them are specially popular: *IntAct* contains binary and complex interactions extracted from the literature [3], *BioGRID* contains protein and genetic interactions derived from reported studies and curated according to the literature [4], finally *Reactome* contains information on chemical reactions involving proteins [5], and uses a particular exhaustive data scheme that allows for the storage of reactions of very different complexities. On the other hand, from protein databases it is possible to retrieve semantic annotations made by researchers. This semantic information could include data on where a protein can be found, what biochemical function it performs or in what biological process it is involved. These annotations used to be written in natural language so it was

difficult to analyze them in a programmatic way. This yielded to further efforts to organize this semantic annotations, lead by the Gene Ontology (GO) project [6]. GO provides a common framework for researchers to annotate proteins and genes with predefined semantic terms. An ontology is a controlled set of terms and relations among them. The *isA* relationship establishes a hierarchy in which a child term refines the meaning of its ancestor term. A parent term can be seen as a generalization of the meaning of all its child terms. GO organizes its terms in three ontologies, each of which contains a different sort of semantic information about a gene or a protein. The *Cellular component* ontology defines possible locations of proteins or expression of genes (e.g. *Cellular membrane* or *Nucleus*). The *Molecular function* ontology defines biochemical functions that a protein, or potentially any molecule, can fulfill (e.g. *Catalytic activity* or *Transporter activity*). Finally, the *Biological process* ontology defines cell processes in which a gene or a protein can be involved (e.g. *Blood coagulation* or *Signal transduction*). The Gene Ontology Annotation (GOA) is a database that contains protein to GO terms annotations[7]. Resnik applied information theory concepts to ontologies [8] defining a semantic similarity using the information content (IC) measure and the *most informative common ancestor* between two terms in an ontology, which have been further extended by other authors (e.g. Jiang and Conrath [9]). The measures cited above define similarity between two terms of an ontology based on the appearance frequency of every term in a knowledge corpus. However, proteins and genes tend to be annotated to sets of terms. Thus, a definition of similarity between sets of terms of an ontology was needed. The Gene Ontology, seen from the *isA* relationship point of view can be seen as a directed acyclic graph (DAG) where nodes are terms and edges go from parents to children nodes. Recently, new measures have been proposed to compare sets of terms of an ontology based on the graph structure induced by them. Falcon and Gentleman proposed an union-intersection based measure called *simUI*, which counts the nodes on the intersection of the two induced graphs and divides it by the number of nodes on the union [10]. This approach was used by Pesquita et al to define *simGIC*, which computes the IC sum of the nodes in the intersection and normalizes it to the IC sum of the nodes in the union [11]. Chagoyen et al used previously proposed measures to define the concept of *coherence* of a protein set [12]. In addition, Ovaska

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et al used ontology-based semantic similarity measures for clustering genes resulting from microarray experiments [13]. According to the authors, this methodology should simplify the analysis of the large number of genes yielded by the experiment. Other authors have employed a similar approach to the prediction of disease candidate genes by means of PPI information [14]. When studying possible genes related to a phenotype, researchers often select target genes based on *a priori* knowledge of gene functions. This can be challenging when the number of candidate genes is very large. Semantic similarity offers a fast and automatic way of comparing *a priori* knowledge of genes.

This manuscript proposes a methodology in order to obtain a set of genes that are candidate to be related with a given phenotype. Given this specific phenotype, the set of already known proteins related to this trait are found. The common set of proteins related with this initial set is obtained through mining of PPI databases. This total set of proteins is organized through a semantic measure and a clustering algorithm in order to obtain a subset of candidates.

II. MATERIALS AND METHODS

The methodology has been applied to two genes known to be related with *Brugada* syndrome phenotype: *SCN5A* and *CACNA1C*. Fig.1 shows further details on the *SCN5A* gene. Defects in *SCN5A* have been related to *Brugada syndrome type* 1 (*BRS1*) [MIM:#601144] and defects in *CACNA1C* have shown relation with *Brugada syndrome type 3 (BRS3)* [MIM:#611875]. Starting from a known set of genes related with a phenotype, the minimum connected subset of the human interactome that contains this set is generated. This is done by growing the map of interactions with new interactants at every iteration until there is only one fully connected component in the graph. Semantic annotations were retrieved for all nodes of the graph, and the extended set of annotations was computed. Extending the set of annotations reflects the fact that if a protein is annotated to a term t then it must also be annotated to all ancestors of t . Then, semantic similarity was computed between all pairs of nodes. To do so IC measure was computed , $IC(t) = -log p(t)$, where $p(t)$ is the probability of appearance of term t in the extended set of annotations[8]. Semantic similarity was computed using *simGIC* measure [11]:

$$
simGIC(p_1, p_2) = \frac{\sum_{t \in \{ann(p_1) \cap ann(p_2)\}} IC(t)}{\sum_{t \in \{ann(p_1) \cup ann(p_2)\}} IC(t)} \tag{1}
$$

where $ann(p)$ is the expanded set of semantic annotations for gene p . From (1) it is obvious that semantic similarity is normalized to 1 for any values of information content. For this reason, IC has not been normalized. Semantic dissimilarity is then computed as:

$$
distGIC(p_1, p_2) = 1 - simGIC(p_1, p_2)
$$
 (2)

Semantic similarity defines a kernel in which genes organize. The goal is to retrieve other genes which are within the minimum semantic hypersphere, given that kernel, that includes all initial genes. The *pam*[15] clustering algorithm

Fig. 1. Description of protein produced by gene SCN5A.

Fig. 2. Graphical description of the proposed methodology.

is employed to partition the data into semantically similar groups. The number of clusters k is defined as the maximum k so that all initial genes are contained in the same cluster. Thus, partitioning the data into $k+1$ groups split the group of initial genes apart (e.g. *SCN5A* and *CACNA1C*). This yielded a set of semantically coherent genes. The overall process is graphically described in Fig. 2.

III. RESULTS

Coronary diseases are polygenic and highly dependent on many environmental factors. The SCN5A gene is part of a sodium channel which mediates the voltage-dependent sodium permeability of excitable membranes. When it is de-

PROTEINS THAT BELONG TO THE SAME CLUSTER AS THE SET OF INPUT PROTEINS.

fective, it modifies the normal Electrocardiogram (ECG) and for some individuals could cause syncope and sudden death. On the other hand, mutations in subunits of calcium channel (CACNA1C) have also been reported for Brugada syndrome related phenotypes. Table I shows the proteins included in the same cluster than the previous proteins, which are able to modify the ECG. These proteins are the most semantically similar to original set (*SCN5A* and *CACNA1C*) in an interaction network. Most proteins in the table are related with voltage dependent calcium channels which are necessary for synaptic transmission. The exocytosis of neurotransmitter vesicles is produced by calcium entrance. Moreover, once the transmission has finished, cell homeostatic equilibrium must be recovered releasing the excess of cations. Any defect in neurotransmission regulation could be the cause of lacks of coordination, cerebral disorders or metabolic problems. Because of this, it is a process strongly controlled in all the stages of cell regulation. From the set of proteins selected *Q9Y2W7* (*Calsenilin*) modulates the transcription of proteins by binding elements of genes in response to calcium concentrations in cells. *Calsenilin* modifies channels density, inactivation kinetics and rate of recovery from inactivation. Other proteins in Table I are directly part of cation channels and some of them have been previously reported to be related with neuromuscular diseases. Calcium channels *Q14643* and *O00555* have been found to be associated with *ataxia* (i.e. lack of coordination of muscle movements). Furthermore, defects in genes coding for *O00305* and *P28472* are associated with neurologic symptoms such as *epilepsy*, *insomnia* and *familiar hemiplegic migraine*, respectively. Ion channels are proteins in membrane which allow the pass of ions in order to maintain a charge concentration gradient between the intracellular and extracellular space. The current conduction generates the action potential necessary for excitable cell functions. An increase of calcium in cytosol is necessary for muscular contraction. Alterations in *P21817* functions interfere in correct sarcoplasmatic reticulum calcium release and in the T-tubules depolarization necessary for muscle contraction. Different myopathies have been related with mutations in this gen. In addition, other proteins connected with the permeability for ions (*P48050*, *P21817* or *Q9UHC3*) are necessary for many processes dependent on osmotic equilibrium. The overall set of obtained proteins is therefore coherent from a biological point of view.

From a computational perspective, obtaining the minimum connecting graph requires checking at every iteration whether the graph is connected. This can be done in time $O(|V| +$ $|E|$, where V is the set of nodes and E is the set of edges. However, for very large graphs it is time consuming. In our case $|V| = 665$ and $|E| = 858$. Calculating semantic similarity between all pairs of genes is the most time and memory consuming task, where $\frac{|V|^2}{2}$ $\frac{1}{2}$ values have to be computed. Each calculation involves two searches in a precalculated annotation list, several searches in a pre-calculated IC table, two sums — the intersection and the union — and a division. A search in the annotation list can be done in the worst case in $O(|V|)$. Each search in the IC table will take $O(|T|)$ where T is the set of all different semantic terms. The number of searches will depend on the average number of annotations per gene, a . Thus, the final cost of this step is:

$$
cost = O(\frac{|V|^2}{2} \cdot |V| \cdot a \cdot |T|) = O(|V|^3 \cdot a \cdot |T|)
$$
 (3)

considering constant sum and division time. Although polynomial, this cost is very non-linear, and it can be very time expensive for high values of |V| and |T|. In this study, $a \simeq 14$ and $|T| = 2157$. Finally, if the starting set of genes is semantically very heterogeneous, calculating the semantic hypersphere that contains them might yield a very large portion of the genes in the graph, which would not be of much interest.

IV. CONCLUSIONS AND FUTURE WORK

From two genes related with a hereditary anomaly that affects heart contraction and cardiac conduction (altering the patient Electrocardiogram), the proposed method is able to obtain a set of genes that have a clear relation with synaptic transmition and muscle contraction. Moreover, it has been reported that mutations in some of them produce different physiological disorders related with these physiological processes. The method is based on mining proteinprotein interaction databases and organizing the set of related proteins through a semantic measure defined from their GO terms. The proposed methodology provides for an automatic way of mining existing curated literature and exploiting it for phenotype related candidate genes generation. This method can effectively be used for finding new candidates related to a disease but also for confirming candidate genes obtained from experimental data, (e.g. microarray datasets of association or linkage analysis).

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¹Rousseeuw et al provided the S original which has been ported to R by Kurt Hornik and has since been enhanced by Martin Maechler: speed improvements, silhouette() functionality, bug fixes, etc. See the 'Changelog' file (in the package source)